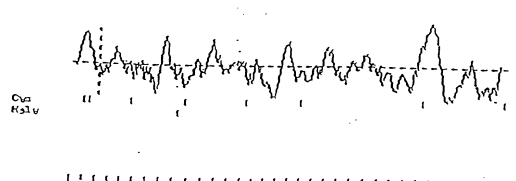
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v	Ħ		S	R		3	t.	C	CGG S. TCA	***	TCA	CYG	ccc	GG	rc	AG (	GAT ·	GGA	GGC	CGA	CAG	GAT	1028 340 1055
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# FIGURE 1a

CCGCTGAACCCCAGGCACCGTATCCAAGGGAGGGATCATGGGAGGGA	1134
CGTGGAAGCAGGAGGCCAGAGGCTACAGCTGTGGAAACGAGGCCATGCTGCTCCTCCTGGTGTTCCATCAGGGAGCCG	1213
TTCGGCCAGTGTCTGTCTGTCTGTCTGCCTCTGTCTGAGGGCACCCTCCATTTGGGATGGAAGGAA	1792
CCCATCCTCCTCCTGCACACTGTGGATGACATGGTACCCTGGCTGG	1371
ANTATGGGCTCCAGACGGATCTCTAAGGTTCCCAGCTCTCAGGGTTGACTCTGTTCCATCCTCTGTGCAAAATCCTCCT	1450
GIGCTICCCTTTGGCCCTCTGTGCTCTTGTCTGGTTTTCCCCAGAAACTCTCACCCTCACTCCCACTGCGGTC	1529
PARCARATETECTTTCGTCTCTCAGAACGGGTCTTGCAGGCAGTTTGGGTATGTCATTCAT	.1608
AGEACGTTGCCCGCTTCCCCTTCACATTAGAAAACAAGATCAGCCTGTGCAACATGGTGAAACCTCATCTCTACCAACAA	1687
WCXXXXXXCXCXXXXTTXGCCXGGTGTGGTGGTGCXTCCCTXTXCTCCCXGCXXCTCGGGGGGCTGAGGTGGGAGA	1766
TGGCTTGAGCCTGGGAGGCAGAGGTTGCAGTGAGCTGAGATCACACCACTGCACTCTAGCTCGGGTGACGAAGCCTGA	1845
CTTGTCTCAAAAAAAACAGGGATGAATATGTCAATTACCCTGATTTGATCATAGCACGTTGTATACATGTACTGCAAT	1924
TTGCTGTCCACCCCATAAATATGTACAATTATGTATACATTTTTAAAATCATAAAAAAATAAGATAATGAAAAAAAA	2003
NNNNNNNNAGGGGGGGCGCTAGACTAGTCTAGAGAACA	2047

# FIGURE 1b

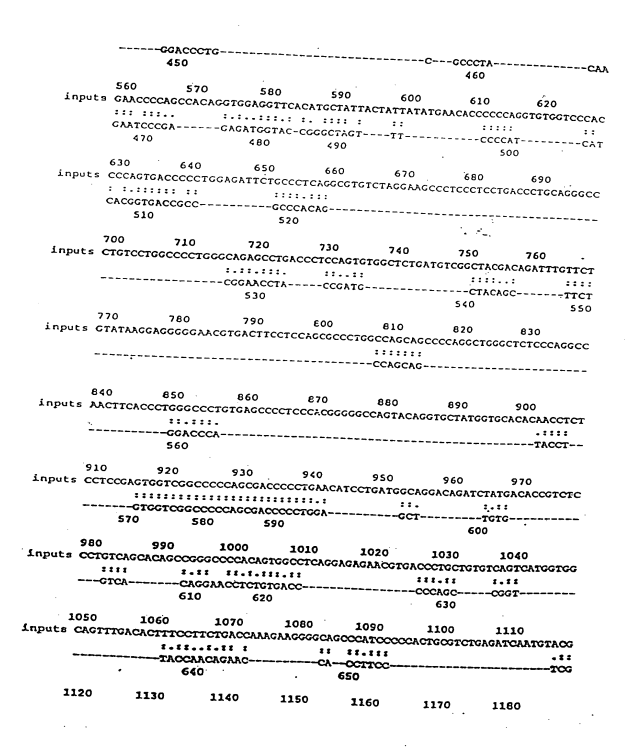


1 41 81 121 161 201 241 281 321

MSPSPTALFCLGLCLGRVPAQSGPLPKPSLQALPSSLVPLEKPVTLRCQGPPGVDLYRLE KLSSSRYQDQAVLFIPAMKRSLAGRYRCSYQNGSLWSLPSDQLELVATGVFAKPSLSAQP GPAVSSGGDVTLQCQTRYGFDQFALYKEGDPAPYKNPERWYRASFPIITVTAAHSGTYRC YSFSSRDPYLWSAPSDPLELVVTGTSVTPSRLPTEPPSSVAEFSEATAELTVSFTNKVFT TETSRSITTSPKESDSPAGPARQYYTKGNLVRICLGAVILIILAGFLAEDWHSRRKRLRH RGRAVQRPLPPLPPLPQTRKSHGGQDGGRQDVHSRGLCS

ALIGN calculates a global alignment of two sequences version 2. OuPlease cite: Hyers and Hiller, CABIOS (1989) > U91928 ORF > hT268 ORF 1893 aa vs. scoring matrix: pam120.mat, gap penalties: -12/-4 37.71 Identity; Global alignment score: -8 . 50 20 30 enter de la companie ATGTCTCCATCCCGACCGCCCTCTTCTGTCTTGGGGCTGTGTCTGGGGGCG-TGTGCCAGC--GCAGAGTG 40 50 90 100 120 inputs GGCCCTTCCCCAAACCCACCCTCTGGGCTGAGCCAGGCTCTGTGAT-CAGCTGGGGGAGCCCCGTGACCA GACCGCTCCCCAAGCCCTCCAGGCTCTGCCCAGCTCCCTGGTGCCCCTGGAGAAGCCA-GTGACCC 110 120 150 160 170 inputs TCTGGTGTCAGGGGAGCCTGGAGGCCCAGGGGTACCGACTGGATAAAGAGGGGAAGCCCAGAGCCCTTGGA 180 TCCGGTGCCAGGG--ACCT-----CCGGGCGTG--GACCTGTA------CCGCCTGGAG-----AAG 160 . 170 210 220 230 240 250 CTGAGTT--CCAGCAGGTACC-AGGATCA-GGCAGTCCTCTTCATCCCGGCCATGAAGAGAAGTCTGGCT 210 220 230 240 280 290 300 310 inputs GGGAGATACCGCTGCCACTATTACAGCTCTGCAG--GCTGGTCAGAGCCCAGCGACCCCCTGGAGCTGGT 320 GGACGCTACCGCTGCTCCTAC--CAGAACGGAAGCCTCTGGTCCCTGCCCAGCGACCAGCTGGAGCTCGT 250 280 290 300 360 370 380 Inputs GATGACAGGATTCTACAACAAACCCACCCTCTCAGCCCTGCCCAGCCCTGTGGTCGCCTCAGGGGGGAAT 390 - 11.121 1 1. - 111111.1 11111111111.111.1111 1 1111 1 11111.111.1 350 360 380 420 430 440 inputs Atcaccroccatgregetcacagaacccatatcaccattttcttcttcatgaaccagaacaccagc 450 elettit teeti - eletetet it tet tittetti tittee tittitt 410 420 . 430 500 510 520 inputs Teccessacetesacteacasetecacastescassessestescassescenterects 530 \*\*\*\*\*\*\* .....

#### FIGURE 3a



#### FIGURE 3b

:		- CIOCICAA	TICCCCXTC.	AGTCCTGTG	ACCTCAGOO	CACGCGGG	
CTA		:::::	***	:. AGAR	11.:	CACUCUGOO,	ACCTACAGG:
660	•	GCAGAA	TTCTC	AGAAC	CCAC		::.:
			670		680	CGCIGA	ACTG
1190							690
inputs CTA	1200	1210	1220	123	0		
·	-OCCTCXTXCX	GCTCC\.k.CCC	CCACCTGCT	STCTTTCCC	12	<0 <sub>1</sub>	250
inputs CTAC : : CC		: :::		• • • • • •	CACTGAGCC	CCTGGAACT	CATGGTCTC
CC	CTCTCATTCA-	Cネ-ネ^C		TCTTCA	•	::::::	
	700		710	YOTTCAC	CAX	-CTGAGACT	TCT-
					-		730
1260	1270 CTCTGGAGGC	1280	1200				
Inputs GCACA						.0 12	120
inputs GCACA	::::	••••••	CYCCCYCYC	GCCCCCCT	CCACACCTG	GTCTCCC	20
	AGGAGTA1	TC 10010-		• • •		- C1000 AA	GATACCTGG
	740	C-ACCACC	y Ciccytyco	GAGTCAG	CTCTCCAC		
	740	,	750	760	770	0166	
1330	1240						•
inputs Acces-	1340 FTC: mm =	1350	1360	1370		_	
inputs ACCTTT	LOSSTTAGEST	CTCGGTGGCC	TTCGTCCTG	CIGCICTT	1380	139	30
			::::		CICCTCCTC	TTCCTCCTC	CTCCGACG
			TCCTC	· ^	:: ::.	:. ::	
			780	C	CCGCCAG	TACTA	22442242
			720		79	0	800
1400	1410	1420	• • • •				
inputs TCAGCG1	CACACCALAC	`ACACCACA	1430	1440	1450	146	•
inputs TCAGCG1		-VCHOOMCY TO	CACCAGAG	AAAGACTGA	TTTCCAGC	1400	, 
CC115		• • • • •					
UCAAC				: ::	:	, recreeked	ACCUTOCC.
OCAAC		-CTGGTC		::: CGGA	:.: :: TAT COO	:: ::	iii:
GCAAC		-CTGGTC 810		CGGA	TATGCC	:: :: TCGG	GGCTG
		ETCCTC		CGGA	TATGCC 820	:: :: TCGG	GGCTG 830
1470	1400	-CTCCTC 810		CGGA	TATGCC	:: :: TCGG	666CTG 830
1470 inputs GAGACAGA	1480 AGCCCAAGGAC	-CTGGTC 810 1490	1500	CGGA	TATGCC	:: :: TCGG	666CTG 830
1470 inputs GAGACAGA	1480 AGCCCAAGGAC	-CTGGTC 810 1490 CAGGGGCCTGG	1500 CTGAGGAGGT	1510 CCAGCCCAG	TATGCC B20 1520	:: :: TCGG	GGCTG 830
1470 inputs GAGACAGA	1480 AGCCCAAGGAC	-CTGGTC 810 1490 CAGGGGCCTGG	1500 CTGAGGAGGT	1510 CCAGCCCAG	TATGCC B20 1520	:: :: TCGG	GGCTG 830
1470 inputs GAGACAGA	1480 AGCCCAAGGAC	-CTGGTC 810 1490 CAGGGGCCTGG	1500 TGAGGAGGT ::::::	1510 CCAGCCAG	TATGCC B20 1520	:: :: TCGG	GGCTG 830
1470 inputs GAGACAGA .::	1480 AGCCCAAGGAC : :: :: :TCCTAATAA- 840	-CTGGTC 810 1490 CAGGGGCCTGG ::::	1500 TGAGGAGGT :::::::	1510 CCAGCCCAG ::: TCTG	TATGCC B20 1520 CCTGCTGACC	:: :: TCGG	GGCTG 830
1470 inputs GAGACAGATGA	1480 AGCCCAAGGAC :::::: TCCTAATAA- 840	-CTGGTC 810 1490 CAGGGGCCTGC ::::	1500 TGAGGAGGT :::::: GCGGGGTT 50	1510 CCAGCCCAG TCTG	1520 SCTGCTGACC	1530 TCCAGGAAG TCCAGGAAG TTCCAGGAAG	GAAAACC
1470 inputs GAGACAGATGA	1480 AGCCCAAGGAC :::::: TCCTAATAA- 840	-CTGGTC 810 1490 CAGGGGCCTGC ::::	1500 TGAGGAGGT :::::: GCGGGGTT 50	1510 CCAGCCCAG TCTG	1520 SCTGCTGACC	1530 TCCAGGAAG TCCAGGAAG TTCCAGGAAG	GAAAACC
1470 inputs GAGACAGATGA 1540 inputs TCTATGCTG	1480 AGCCCAAGGAC IIIIIIIIIIIIIIIIIIIIIIIIIII	-CTGGTC 810  1,490 CAGGGGCCTGGTCCTG 8  1560	1500 TGAGGAGGI :::::: GCGGGGTT 50 1570 IGAGG-ACAG	1510 CCAGCCCAG TCTG 860 1580	1520 CTGCTGACG ::-::::::::::::::::::::::::::::::::::	1530  TCCAGGAA  TSTCCAGGAA  TSTCCAGAA  TSTCCAGAA  TSTCCAGAAA  TSTCCAGAAA  TSTCCAGAAAA  TSTCCAGAAAA  TSTCCAGAAAA  TSTCCAGAAAAA  TSTCCAGAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA	GAAAACC  GAAAACC  GAAAACC
1470 inputs GAGACAGATGA 1540 inputs TCTATGCTG	1480 AGCCCAAGGAC IIIIIIIIIIIIIIIIIIIIIIIIIII	-CTGGTC 810  1,490 CAGGGGCCTGGTCCTG 8  1560	1500 TGAGGAGGI :::::: GCGGGGTT 50 1570 IGAGG-ACAG	1510 CCAGCCCAG TCTG 860 1580	1520 CTGCTGACG ::-::::::::::::::::::::::::::::::::::	1530  TCCAGGAA  TSTCCAGGAA  TSTCCAGAA  TSTCCAGAA  TSTCCAGAAA  TSTCCAGAAA  TSTCCAGAAAA  TSTCCAGAAAA  TSTCCAGAAAA  TSTCCAGAAAAA  TSTCCAGAAAAAAAAA  TSTCCAGAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA	GAAAACC  GAAAACC  GAAAACC
1470 inputs GAGACAGATGA 1540 inputs TCTATGCTG	1480 AGCCCAAGGAC IIIIIIIIIIIIIIIIIIIIIIIIIII	-CTGGTC 810  1,490 CAGGGGCCTGGTCCTG 8  1560	1500 TGAGGAGGI :::::: GCGGGGTT 50 1570 IGAGG-ACAG	1510 CCAGCCCAG TCTG 860 1580	1520 CTGCTGACG ::-::::::::::::::::::::::::::::::::::	1530  TCCAGGAA  TSTCCAGGAA  TSTCCAGAA  TSTCCAGAA  TSTCCAGAAA  TSTCCAGAAA  TSTCCAGAAAA  TSTCCAGAAAA  TSTCCAGAAAA  TSTCCAGAAAAA  TSTCCAGAAAAAAAAA  TSTCCAGAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA	GAAAACC  GAAAACC  GAAAACC
1470 inputs GAGACAGATGA 1540 inputs TCTATGCTG	1480 AGCCCAAGGAC :::::: ATCCTAATAA - 840 1550 GCCGTGAAGG	-CTGGTC 810  1,490 CAGGGGCCTGGTCCTG 8  1560	1500 TGAGGAGGI :::::: GCGGGGTT 50 1570 IGAGG-ACAG	1510 CCAGCCCAG TCTG 860 1580 GGGTGGAGCC	1520 CCTGCTGACG :::::::::::::::::::::::::::::::::::	1530 FTCCAGGAA FTCCAGAGGCCCA FTCCAGAGGCCCA FTCCAGAGGCCCA	GAAAACC  GAAAACC  GAAAACC
1470 inputs GAGACAGATGA  1540 inputs TCTATGCTCAG	1480 AGCCCAAGGAC :::::::: ATCCTAATAA- 840 1550 GCCGTGAAGGA	1490 CAGGGCCTGC :::TCCTG	1500 CTGAGGAGGT ::::::GCGGGGTT 50 1570 IGAGG-ACAG	1510 CCAGCCCAG ::: TCTG 860 1580 GGGTGGAGCC	1520 CTGCTGACC ::::::::GCAGA-C	1530  TCCAGGAA  TSTCCAGGAA  TSTCCAGAA  TSTCCAGAA  TSTCCAGAAA  TSTCCAGAAA  TSTCCAGAAAA  TSTCCAGAAAA  TSTCCAGAAAA  TSTCCAGAAAAA  TSTCCAGAAAAAAAAA  TSTCCAGAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA	GAAAACC  GAAAACC  GAAAACC
1470 inputs GAGACAGETGA  1540 inputs TCTATGCTG ACAG	1480 AGCCCAAGGAC ::::::: ATCCTAATAA- 840 1550 GCCGTGAAGGA	-CTGGTC 810  1490 CAGGGGCCTGG ::::TCCTG- 8  1560 ACACACACACTCT ::: LAGGGGCT	1500 CTGAGGAGGT :::::::CCGGGGTT 50 1570 IGAGG-ACAG	1510 CCAGCCCAG :: TCTG 860 1580 GGGTGGAGCC	1520 CCTGCTGACC :::::::::::::::::::::::::::::::::::	1530 GTCCAGGAAG : ::: GGACTGG 870 160 CAGAGCCCA	GAAAACC GAAAACCC OO.CAOGAT
1470 inputs GAGACAGE -::TGA  1540 inputs TCTATGCTG ACAG	1480 AGCCCAAGGAC ::::::: ATCCTAATAA- 840 1550 GCCGTGAAGGA	-CTGGTC 810  1,490 CAGGGGCCTGC ::::TCCTG 8  1560 ACACACAGTCT :::::::: LAGGGCCT 890  1630 CCTATGCCCCC	1500 TGAGGAGGT ::::::CCGGGGTT 50 1570 IGAGG-ACAG :::::::::: IGCGGGACAG 900 1640 GGIGAAAGA	1510 CCAGCCCAG :: TCTG 860 1580 GGGTGGAGCC	1520 CCTGCTGACC :::::::::::::::::::::::::::::::::::	1530 GTCCAGGAAG : ::: GGACTGG 870 160 CAGAGCCCA	GAAAACC GAAAACC  GAAAACC  GAAAACC  GAAAACC  GAAAACC  GAAAACC  GAAAACC  GAAAACC  GAAAACC  GAAAACC  GAAAACC  GAAAACC  GAAAACC  GAAAACC  GAAAAACC  GAACAACAACAACAACAACAACAACAACAACAACAACAAC
1470 inputs GAGACACACACACACACACACACACACACACACACACAC	1480 AGCCCAAGGAC ::::::: ATCCTAATAA- 840 1550 GCCGTGAAGGA :::::::::: GCCGGAGGA 880 1620 CAGGCAGTGA	-CTGGTC 810  1490 CAGGGGCCTGG :::TCCTG- 8  1560 ACACACACACTCT ::: LAGGGCCT 890  1630 CCTATGCCCC	1500 CTGAGGAGGT :::::::CCGGGGTT 50 1570 IGAGG-ACAG IIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII	1510 CCAGCCCAG :: TCTG 860 1580 GGGTGGAGCT ::: GGGCA	1520 CCTGCTGACC :::::::::::::::::::::::::::::::::::	1530 GTCCAGGAAG : ::: GGACTGG 870  160 CAGAGCCCA ::::: CAGAGCCCA 920	GAAAACC  GAAAACC  GAAAACC  GAAAACC  GACGAT  GA
1470 inputs GAGACACACACACACACACACACACACACACACACACAC	1480 AGCCCAAGGAC ::::::: ATCCTAATAA- 840 1550 GCCGTGAAGGA :::::::::: GCCGGAGGA 880 1620 CAGGCAGTGA	-CTGGTC 810  1490 CAGGGGCCTGG :::TCCTG- 8  1560 ACACACACACTCT ::: LAGGGCCT 890  1630 CCTATGCCCC	1500 CTGAGGAGGT :::::::CCGGGGTT 50 1570 IGAGG-ACAG IIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII	1510 CCAGCCCAG :: TCTG 860 1580 GGGTGGAGCT ::: GGGCA	1520 CCTGCTGACC :::::::::::::::::::::::::::::::::::	1530 STCCAGGAAG STCCAGGAAG STCCAGGAAG STCCAGGAAG STCCAGGAAG STCCAGGAAG STCCAGAGGCCCAGAAGGCCCAGAAGGCCCAGAAGGCCCAGAAGGCCCAGAAGGCCCAGAAGGCCCAGAAAGGCCCAGAAAGGCCCAGAAAGGCCCAGAAAGGCCCACAAAAGGCCCACAAAAGGCCCCACAAAAGGCCCCACAAAAGGCCCCACAAAAGGCCCCACAAAAGGCCCCACAAAAGGCCCCACAAAAGGCCCCACAAAAAGGCCCCACAAAAAGGCCCCACAAAAAA	GAAAACC  GAAAACC  COCOCOCOCOCOCOCOCOCOCOCOCOCOCOCO
1470 inputs GAGACACACACACACACACACACACACACACACACACAC	1480 AGCCCAAGGAC ::::::: ATCCTAATAA- 840 1550 GCCGTGAAGGA :::::::::: GCCGGAGGA 880 1620 CAGGCAGTGA	-CTGGTC 810  1490 CAGGGGCCTGG :::TCCTG- 8  1560 ACACACACACTCT ::: LAGGGCCT 890  1630 CCTATGCCCC	1500 CTGAGGAGGT :::::::CCGGGGTT 50 1570 IGAGG-ACAG IIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII	1510 CCAGCCCAG :: TCTG 860 1580 GGGTGGAGCT ::: GGGCA	1520 CCTGCTGACC :::::::::::::::::::::::::::::::::::	1530 STCCAGGAAG STCCAGGAAG STCCAGGAAG STCCAGGAAG STCCAGGAAG STCCAGGAAG STCCAGAGGCCCAGAAGGCCCAGAAGGCCCAGAAGGCCCAGAAGGCCCAGAAGGCCCAGAAGGCCCAGAAAGGCCCAGAAAGGCCCAGAAAGGCCCAGAAAGGCCCACAAAAGGCCCACAAAAGGCCCCACAAAAGGCCCCACAAAAGGCCCCACAAAAGGCCCCACAAAAGGCCCCACAAAAGGCCCCACAAAAGGCCCCACAAAAAGGCCCCACAAAAAGGCCCCACAAAAAA	GAAAACC  GAAAACC  COCOCOCOCOCOCOCOCOCOCOCOCOCOCOCO
1470 inputs GAGACAGA  1540 inputs TCTATGCTG  ACAG  1610 Inputs GAAGACCCCC TCC	1480 AGCCCAAGGAC IIIIIIIIIIIIIIIIIIIIIIIIIII	-CTGGTC 810  1,490  CAGGGGCCTGC ::::TCCTG 8  1560  ACACACAGTCT :::: 1,460GCCT 890  1630  CCTATGCCCCC ::::: 930	1500 CTGAGGAGGT :::::::CCGGGGTT 50 1570 IGAGG-ACAG IIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII	1510 CCAGCCCAG :: TCTG 860 1580 GGGTGGAGCT ::: GGGCA	1520 CCTGCTGACC :::::::::::::::::::::::::::::::::::	1530 GTCCAGGAAG FILE GGACTGG 870  CAGAGCCCA FILE CAGAGCCCA FILE CAGAGCCCC FILE FILE CAGAGCCCC FILE CAGAGCCCC FILE FILE CAGAGCCCC FILE CAGAGCCCC FILE CAGAGCCCCC FILE CAGAGCCCCC FILE CAGAGCCCCC FILE CAGAGCCCCCC FILE CAGAGCCCCCC FILE CAGAGCCCCCC FILE CAGAGCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	GAAAACC  GAAAACC  CACGAT  CCT  CTCCTC  CCTCCTC
1470 inputs GAGACAGA .::TGA  1540 inputs TCTATGCTG .: ACAG 1610	1480 AGCCCAAGGAC :::::::::::::::::::::::::::	-CTGGTC 810  1,490 CAGGGGCCTGC ::::TCCTG 8  1560 ACACACACTCT :::: LAGCGCCT 890  1630 CGTATGCCCCC :::::GCCCCT 930	1500 CTGAGGAGGT :::::: -GCGGGGTT 50 1570 IGAGG-ACAG II :::::: IGCGGGACAG 900 1640 GGTGAAACAG	1510 CCAGCCCAG ::: TCTG 860 1580 GGGTGGAGC: ::: GGGGCA	1520 CTGCTGACC ::::: GCAGA-C 1590 IGGACAGT- :::::: GGGCTGTG 910 1660 TAGGAGAGA	1530 GTCCAGGAAG FILE GGACTGG 870  160 CAGAGCCCA FILE CAGAGCCC 920  1670 AATGGCCT 940	GAAAACC  GAAAACC  CO CAOGAT  CO CTCCTC  CTCCTC
1470 inputs GAGACAGA .::TGA  1540 inputs TCTATGCTG .: ACAG 1610	1480 AGCCCAAGGAC :::::::::::::::::::::::::::	-CTGGTC 810  1,490 CAGGGGCCTGC ::::TCCTG 8  1560 ACACACACTCT :::: LAGCGCCT 890  1630 CGTATGCCCCC :::::GCCCCT 930	1500 CTGAGGAGGT :::::: -GCGGGGTT 50 1570 IGAGG-ACAG II :::::: IGCGGGACAG 900 1640 GGTGAAACAG	1510 CCAGCCCAG ::: TCTG 860 1580 GGGTGGAGC: ::: GGGGCA	1520 CTGCTGACC ::::: GCAGA-C 1590 IGGACAGT- :::::: GGGCTGTG 910 1660 TAGGAGAGA	1530 GTCCAGGAAG FILE GGACTGG 870  160 CAGAGCCCA FILE CAGAGCCC 920  1670 AATGGCCT 940	GAAAACC  GAAAACC  CO CAOGAT  CO CTCCTC  CTCCTC
1470 inputs GAGACAGA .::TGA  1540 inputs TCTATGCTG .: ACAG 1610	1480 AGCCCAAGGAC :::::::::::::::::::::::::::	-CTGGTC 810  1.490 CAGGGGCCTGG :::TCCTG 8  1560 ACACACACACTCT ::: AGCGCCT 890  1630 CCTATGCCCCC 930  1700 TTCCTGCACA	1500 CTGAGGAGGT ::::::CCCGGGTT 50 1570 IGAGG-ACAG ::::::::: IGCGGCACAG 900 1640 GGTGAAACAG : IG 1710 CAAAGGACA	1510 CCAGCCCAG ::: TCTG 860 1580 GGGTGGAGCC ::: FGGGCA 1650 CTCCAGTCC	1520 CTGCTGACC ::::: GCAGA-C 1590 IGGACAGT- :::::: GGGCTGTG 910 1660 TAGGAGAGA	1530 GTCCAGGAAG FILE GGACTGG 870  160 CAGAGCCCA FILE CAGAGCCC 920  1670 AATGGCCT 940	GAAAACC  GAAAACC  CO CAOGAT  CO CTCCTC  CTCCTC
1470 inputs GAGACAGA .::TGA  1540 inputs TCTATGCTC .: ACAC B I610 Inputs GAAGACCCC .::TCC I680 IPUts CCTCCTCACT	1480 AGCCCAAGGAC IIIIIIIIIIIIIIIIIIIIIIIIIII	-CTGGTC 810  1,490 CAGGGGCCTGG ::::TCCTG 8  1560 ACACACAGTCT :::: LAGGGCCT 890  1630 CCTATGCCCGG ::::: 930  1700 TTCCTGGACA	1500 CTGAGGAGGT  ::::::CCGGGGTT  50  1570 IGAGG-ACAG  900  1640 GGTGAAACAG  : IG  1710 CAAAAGGACA	1510 CCAGCCCAG  1510 TCTG 860 1580 GGCTGGAGCC 1550 CTCCAGTCC 1720 GACAGGTGG	1520 CTGCTGACC CTGCTGACC CTGCTGACC CTGCAGACC CTGCACACC CTCACACC CTCACACCC CTCACACCC CTCACACCC CTCACACCC CT	1530 GTCCAGGAAG : ::: GGACTGG 870  160 CAGAGCCCA ::::: CAGAGCCCA ::::: CAGAGCCCA 920  1670 AATGGCCTC 940  1740 AGGCAGATC	GAAAACC  GAAAACC  CACGAT  CT  CTCCTC  CACGAC
1470 inputs GAGACAGA  1540 inputs TCTATGCTC  ACAC  AGACACCCC  TITLE  1680 Aputs CCTCCTCACT  IIIII I.I.	1480 AGCCCAAGGAC IIIIIIIIIIIIIIIIIIIIIIIIIII	-CTGGTC 810  1,490 CAGGGGCCTGG ::::TCCTG 8  1560 ACACACAGTCT :::: LAGGGCCT 890  1630 CCTATGCCCGG ::::: 930  1700 TTCCTGGACA	1500 CTGAGGAGGT  ::::::CCGGGGTT  50  1570 IGAGG-ACAG  900  1640 GGTGAAACAG  : IG  1710 CAAAAGGACA	1510 CCAGCCCAG  1510 TCTG 860 1580 GGCTGGAGCC 1550 CTCCAGTCC 1720 GACAGGTGG	1520 CTGCTGACC CTGCTGACC CTGCTGACC CTGCAGACC CTGCACACC CTCACACC CTCACACCC CTCACACCC CTCACACCC CTCACACCC CT	1530 GTCCAGGAAG : ::: GGACTGG 870  160 CAGAGCCCA ::::: CAGAGCCCA ::::: CAGAGCCCA 920  1670 AATGGCCTC 940  1740 AGGCAGATC	GAAAACC  GAAAACC  CACGAT  CT  CTCCTC  CACGAC
1470 inputs GAGACAGA .::TGA  1540 inputs TCTATGCTC .: ACAC B I610 Inputs GAAGACCCC .::TCC I680 IPUts CCTCCTCACT	1480 AGCCCAAGGAC IIIIIIIIIIIIIIIIIIIIIIIIIII	-CTGGTC 810  1.490 CAGGGGCCTGG :::TCCTG 8  1560 ACACACACACTCT ::: AGCGCCT 890  1630 CCTATGCCCCC 930  1700 TTCCTGCACA	1500 CTGAGGAGGT  ::::::CCGGGGTT  50  1570 IGAGG-ACAG  900  1640 GGTGAAACAG  : IG  1710 CAAAAGGACA	1510 CCAGCCCAG  1510 TCTG 860 1580 GGCTGGAGCC 1550 CTCCAGTCC 1720 GACAGGTGG	1520 CTGCTGACC CTGCTGACC CTGCTGACC CTGCAGACC CTGCACACC CTCACACC CTCACACCC CTCACACCC CTCACACCC CTCACACCC CT	1530 GTCCAGGAAG : ::: GGACTGG 870  160 CAGAGCCCA ::::: CAGAGCCCA ::::: CAGAGCCCA 920  1670 AATGGCCTC 940  1740 AGGCAGATC	GAAAACC  GAAAACC  CO  CACGAT  CO  CTCCTC  CO  GACAC  II

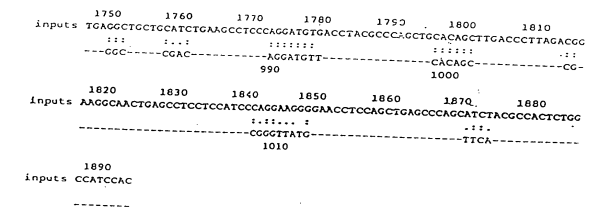
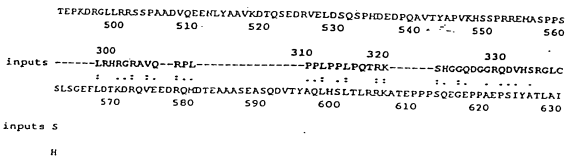


FIGURE 3d

```
ALIGN calculates a global alignment of two sequences
              version 2. Ouplease cite: Hyers and Hiller, CABIOS (1989)
            > hT268 a.a.
           > GenPept U91928 - Human clone HL9 monocyte inhib
           scoring matrix: pam120.mat, gap penalties: -12/-4
           23.01 identity;
                                                                      Global alignment score: -642
                                                                              20
                                                                                                        30
          Inputs HSPSPTALFCLGLCLG-RVPAQSGPLPKPSLQALPSSLVPLEKPVTLRCQGPPGVDLYRLEKLSSS----
                                                                                                                                     40
                                                                                                                                                           . 50
                            tere firming in the experience of the contract of the contract
                            HTPALTALLCLGLSLGPRTRVQAGPFPKPTLHAEPGSVISHGSPVTIHCQGSLEAQEYRLDKEGSPEPLD
                                                                    . 20
                                                                                              30
                                                                                                                             40
                                                                                                                                                            50
                                                70
                                                                                                                                                  110 . -_ 120
                                                                                                    90
                                                                                                                           100
        inputs RYQ-----DQAVLFIPAMKRSLAGRYRCSYQNGSLHSLPSDQLELVATGVFAKPSLSAQPGPAVSSGGDV
                           ti di con la constitución de la 
                           RUMPLEPKNXARFSIPSHTEHHAGRYRCHYYSSAGHSEPSDPLELVHTGFYNKPTLSALPSPVVASGGM
                                                               90 100
                                                                                                                       110 120
      inputs TLQCQT-----RY-----RY-----
                         TLRCGSQKGYHHFVLMKEGEHQLPATLDSQQLHSGGFQALFPVGPVNPSHRWRFTCYYYYMYTPQVWSHP
                                                                  160
                                                                                                                         180 190
                                                                                                                               140
    inputs -----GFDQFALYKEGDP-----
                                                                                                                                : - : - : - : : : : : .
                        SDPLEILPSGVSRKPSLLTLQGPVLAPGQSLTLQGGSDVGYDRFVLYKEGERDFLQRPGQQPQAGLSQAN
                                                       230 240 250
   inputs -----APYK-----ERW--
                      FILGPVSPSHGGQYRCYGAHNLSSEHSAPSDPLNILMAGQIYDTVSLSAQPGPTVASGENVTLLCQSHHQ
                                                                300 310
                                                                                                                320 330 .340
                                                                                                              170
                                                                                                                                         180
                                                                                                                                                                                                     200
                                                                                                                                                                        190
  Inputs ----YRASFPIITVTAAHSGTYRCYSFSSRDPYLWSAPSDPLELVVTG
                                                                                          FOTFLLTKEGAAHPPLRLRSHYGAHKYQAEFPHSPVTSAHAGTYRCYGSYSSHPHLLSFPSEPLELHVSG
                                       360
                                                    370 380 390
                                                    220 230
                                                                                                                                       240
                                                                                                                                                                    250
HSGGSSLPPTGPPSTPGLGRYLEVLIGVSVAFVLLLFLLLFLLLRRQRHSKHRTSDQRKTDFQRPAGAAE
                                                             440
                                                                                    450
                                                                                                          460
                                                                                                                                       470
                                                              280
                                   270
                                                                                                      290
inputs ROYYIKCHLVRICLGAVIL----IILAGFLAEDH-
                                                                                                                                         .::: .:.
                                                                                     :: .:
```

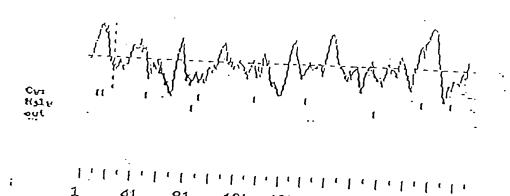
#### FIGURE 4a



••

GAGTCGACCCACGCGTCCCGCTTCCCTGCTTGGCCACATAGCTCAGGACTGGGTTGCAGAACC ATG TCT CCA GCC	7 4
S P T F F C I G L C V L Q V I Q T Q S G TCA CCC ACT. TTC TCT ATT GGG CTG TGT GTA CTG CAA GTG ATC CAA ACA CAG AGT GGC 13	
P L P K P S L Q A Q P S S L V P L G Q S CC C	٠ ،
V I L R C Q G P P D V D L Y R L E K L K 6 GTT ATT CTG AGG TGC CAG GGA CCT CCA GAT GTG GAT TTA TAT CGC CTG GAG AAA CTG AAA 25	<b>.</b>
PEKYED QDFLFIPTHERS NA 8 CCG GAG AAG TAT GAA GAT CAA GAC TIT CTC TTC ATT CCA ACC ATG GAA AGA AGT AAT GCT 31	<
G. R Y R C S Y Q N G S H W S L P S. D Q L 10 GGA CGG TAT CGA TGC TCT TAT CAG ANT GGG AGT CAC TGG TCT CTC CCC 15TT C10	4
E L I A T C V Y A K P S L S A K P S S S A 12.	<
V P Q G R D V T L K C Q S P Y S F D E F 144	(
V L Y K E G D T G P Y K R P E K W Y R A 166	ŧ.
N F P I I T V T A A H S G T Y R C Y S F 186	:
S S S P Y T W S I D S D D D D D D D D D D D D D D D D	
THE AGE TEA TET CEA TAC CTG TGG TEA GEE CEG AGT GAC CET CTA GTG CTT GTG GTT ACT 674  G L S A T P S Q V P T E E S F P V T E S 224	į
SGA CTC TCT GCC ACT CCC AGC CAG GTA CCC ACG GAA GAA TCA TTT CCT GTG ACA GAA TCC 736  S R R P S I L P T H K I S T T E K P H H 246	
TO AGG AGA COT TOO ATO TTA COO ACA AAC AAA ATA TOT ACA ACT GAA AAG COT ATG AAT 794	
TO ACT GCC TCT CCA GAG GGG CTG AGC CCT CCA ATT GGT TTT GCT CAT CAG CAC TAT GCC 854  K G N L V R I C L G A T I I I L L G L 284	
AG GGG AAT CTG GTC CGG ATA TGC CTT GGT GCC ACG ATT ATA ATA ATT TTG TTG GGG CTT 914  L A E D W H S R K K C L Q H R H R A L Q 304	
R P L P P 1. P 1. A 1.	
GG CCA CTA CCA CCC CTC CCA CTG GCC TAG  1004  AATAACTTGGCTTTCAGCAGAGGGATTGACCAGACATCCATGCACCACTGGACATCACCACTAGAGCCACAGACAT _ 108	
CACATACTCAAGAGTGGGGAGGTTATATAAAAAAAATGAGTGAG	
. 116:	3

41

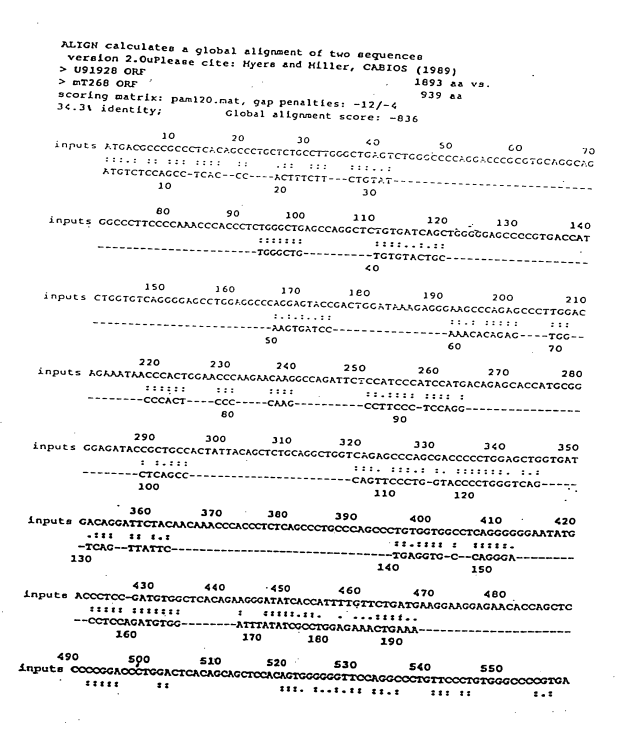


121

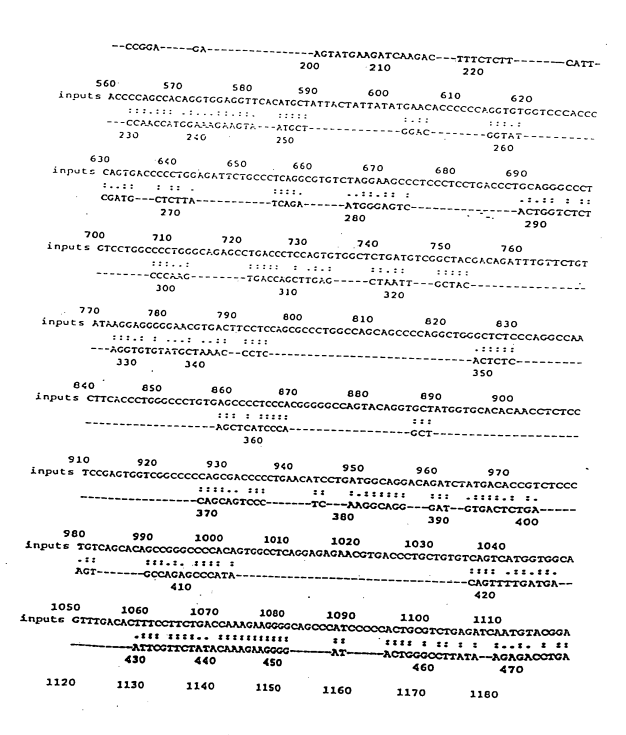
MSPASPTFFCIGLCVLQVIQTQSGPLPKPSLQAQPSSLVPLGQSVILRCQGPPDVDLYRL EKLKPEKYEDQDFLFIPTMERSNAGRYRCSYQNGSHWSLPSDQLELIATGVYAKPSLSAH PSSAVPQGRDVTLKCQSPYSFDEFVLYKEGDTGPYKRPEKWYRANFPIITVTAAHSGTYR CYSFSSSPYLWSAPSDPLVLVVTGLSATPSQVPTEESFPVTESSRRPSILPTNKISTTE KPMNITASPEGLSPPIGFAHQHYAKGNLVRICLGATIIIILLGLLAEDWHSRKKCLQHRM

201

161



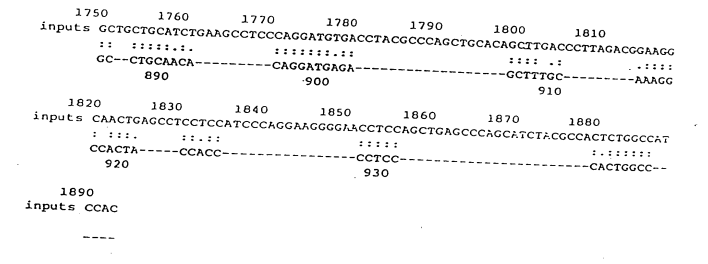
### FIGURE 8a



#### FIGURE 8b

```
Inputs GCTCATAAGTACCAGGCTGAATTCCCCATGAGTCCTGTGACCTCAGCCCACGCGGGGACCTACAGGTGCT
                     G--AAATGGTACCGGGCCAATTTCCCCATCATCACAGTGACTGCTGCTCACAGTGGGACGTACCGGTCTT
                                                                                       520
                                                                                                        530
                               1200
                                              1210
          inputs ACGGCTCATACAGCTCCAACCCCCACCTGCTGTCTTTCCCCAGTGAGCCCCTGGAACTCATGGTCTCAGG
                    ACAGCTTCTCCAGCTCATCTCCATACCTGTGGTCAGCCCCGAGTGACCCTCTAGTGCTTGTGGTTACTGG
                                                                    580 590
               1260
                              1270 1280
        inputs ACACTCTGGAGGCTCCAGCCTCCACACGGGCCGCCTCCACACCTGGTCTGGGAAGATACCTGGAG
                   ACTCTCTG----CCA--CTCCCAGCC--AGGT--ACCCAC-----GGA-AGAATCATTTCCTG---
             1330 1340
                                            1350 1360
       ----TGA-----CAGAATCCT----CCAGGAGACCTTCCA----TCTTAC----CCACAAACAAA
                         1<10
                                     1420 1430
     inputs AGCGTCACAGCAAACACAGGACATCTGACCAGAGAAAGACTGATTTCCAGCGTCCTGCAGGGGCTGCGGA
                i fatari tara antimitana a fataritaria
                A---TATCTACA4---CTGAA----AAGCCTATGAATATC--ACTGCCT-C-TCCAG-AGGGGCTG----
                                                       730
                        1480 1490
    Inputs GACAGAGCCCAAGGACAGGGGCCTGCTGAGGAGGTCCAGGCCCAGCTGCTGACGTCCAGGAAGAAAACCTC
                                                       1500
                      ----AGCCCT-----CC----AATTGGTTTTGCTCATCAGCA-----C
                                                                          780
         1540 1550
  Inputs TATGCTGCCGTGAAGGACACAGTCTGAGGACAGGGTGGAGGTGAGAGTCAGAGGACACGATGAAG
                                       1560
                                                         rer rer reres
             TATGC-----CGGATATG
                                                                        800
                      1620 1630 1640 1650 1660
 Inputs Acceccageagtaacgtatgcccccgtgaaacactccagtcctaggagaaaatggcctctcctcctc
              Y at a state and a
            ---CCTTGG-----TGCCACGAT-----TATAATAATTTTGT-----
                    1690
                                 1700 1710
Inputs CTCACTGTGGGGAATTCCTGGACACAAAGGACAGACAGGTGGAAGAGGACAGGCAGATGGACACTGAG
                      ***** .: ::-: ::-::: .:.:
                 -----TGGGGCTT--CTAG---CAGAGGATTGGC------ACAGTCGGAAGAA---
                                                                             **** ****** *
                                                     860
                                                                                         870
```

## FIGURE 8c



```
ALIGN calculates a global alignment of two sequences
           version 2. OuPlease cite: Hyers and Hiller, CABIOS (1989)
         > mT268 a.a.
         > GenPept U91928 - Human clone HL9 monocyte inhib. 631 aa
                                                                                                 313 aa vs.
         scoring matrix: pam120.mat, gap penalties: -12/-4
         20.31 Identity;
                                               Global alignment score: -802
                                  10
        inputs HSPASPTFFCIGLCVLQVIQTQSGPLPKPSLQAQPSSLVPLGQSVILRCQGPPDVDLYRLEKL-KPZEYE
                                                                   30
                    the contract of the contract o
                    HTPALTALLCLGLSLGPRTRVQAGPFPKPTLHAEPGSVISHGSPVTIHCQGSLEAQEYRLDKEGSPEPLD
                                                             30 40
                                                                                                   50
                                              80
       inputs DQDFL-----F-IPTHERSHAGRYRCSYQNGSHWSLPSDQLELIATGVYAKPSLSAHPSSAVPQGRDV
                                                                              100
                   RNNPLEPKNKARFS I PSHTEHHAGRYRCHYYSSAGHSEPSDPLELVHTGFYNKPTLSALPSPVVASGGI::
                                                                                110 120 130
     inputs TLKC--QSPY-----
                 TLRCGSQKGYHHFVLHKEGEHQLPRTLDSQQLHSGGFQALFPVGPVNPSHRWRFTCYYYYHHTPQVWSHP
                                                                              180
    inputs -----SFDEFVLYKEGO-----
                SDPLEILPSGVSRKPSLLTLQGPVLAPGQSLTLQCGSDVGYDRFVLYKEGERDFLQRPGQQPQAGLSQAM
                                                                                    ... :::::::
                                                      240
                                                                       250
                                                                                           260
   inputs -----RP-----EKW--
               FTLGPV5P5HGGQYRCYGAHNLSSEHSAPSDPLNILHAGQIYDTV5L5AQPGPTVASGENVTLLCQSHXQ
                                           300 310 320 330 340
                                                                     170
                                                                                     180
                                                                                                      190
                                              -----Yranfpiitvtaahsgtyrcysfsssspylhsapsdplvlvvtg
                                                        intette trettettitie itet i i itest feter
              FOTFLLTKEGAAHPPLRLRSHYGARKYQAEFPHSPVTSAHAGTYRCYGSYSSHPRLLSFPSEPLELHVSG
                         360 370
                                                       380
                                                                            390
                                                                                            400
                                                                                                               410
                                                               220
inputs LSATPSQVPTEES-----FPV-----FPV-----
               HSGGSSLPPTGPPSTPGLGRYLEVLIGVSVAFVLLLFLLLFLLLRRQRHSKHRTSDQRKTDFQRPAGAAE
                                         440 450 460 470
                                         230
                                                                240
                                                                                  250
::.:
```

### FIGURE 9a

TEPKDRGLLRRSSPAADVQEENLYAAVKDTQSEDRVELDSQSPHDEDPQAVTYAPVTHSSPRPEMASPPS

500 510 520 530 540 550 560

280 290 300 310

inputs CLGATIIIILLGLLAEDWH-----SRKKCLQHRHRALQRPL----PP------LPL

51.5CEFLDTKDRQVEEDRQHDTEAAASEASQDVTYAQLHSLTLRRKATEPPPSQEGEPPAEPSIYATLAI

570 580 590 600 610 620 630

inputs A

FIGURE 9b

```
ig: domain 2 of 2, from 128 to 183: score 9.6, E = 1.6

*->GesvtLtCsvsgfgppgvsvtWyfkngk.lgpsllgysysrlesgek
G +vtL C++ ++ y k+g++ + y+r+e +
GRDVTLKCQSP---YSFDEFVLY-KEGDtGP-------YKRPEKW-Y 162

anlsegrfsissltLtissvekeDsGtYtCvv<-*
+ ++i++v++ sGtY+C

mT268 163 RA-------NFPIITVTAAHSGTYRCYS 183
```

```
ALIGN calculates a global alignment of two sequences
   version 2.0uPlcase cite: Myers and Hiller, CABIOS (1989)
  > hT268 a.a.
                                          339 ac vs.
  > mT268 a.a.
                                          313 ca
  scoring matrix: pan120.mat, gap penalties: -12/- < K
  64.4% identity;
                Global alignment score: 1011
                      20
                             30
                                     40
 inputs HSPSPTALFCLGLCLGRV-PADSGPLPKPSLQALPSSLVPLEKPVTLRCQGPPGVDLYRLEKLSSSRYQD
                                             50
       HSPASPTFFCIGLCVLQVIQTDSGPLPKPSLQAQPSSLVPLGQSVILRCQGPPDVDLYRLEKLKPEKYED
                     20
                                    <0
                                            50
      70
             03
                     90
                            100
                                    110
                                           120
 inputs QAVLFIPATRESLAGRYRCSYQNGSLHSLPSDQLELVATGVFAKPSLSAQPGPAVSSGGDVTLQCQTRYG
      QDFLF1PTHERSHAGRYRCSYQNGSHWSLPSDQLELIATGVYAKPSLSAMPSSAVPQGRDVTLKCQSPYS
            20
                    90
                           100
                                   110 .
                                          120
                                                  130 .
            150
                    160
                            170
                                   180
                                           190
inputs FDOFALYKEGDPAPYKNPERWYRASFPIITVTAAHSGTYRCYSFSSRDPYLWSAPSDPLELVVTGTSVTP
                                                   200
      FDEFVLYKEGDTGPYKRPEKWYRAITPIITVTAAHSGTYRCYSFSSSSPYLWSAPSDPLVLVVTGLSATP
                   160
                           170
                                  180
                                                  200
                   230
                           240
                                   250
                                           260
inputs splpteppssvaefseataeltvsftnkv77tetsrsittspkesdspagparqyytkgnlvkiclgavi
      SQVPTEESFPVTESSRRPSILP---TNKISTTEKPYNITASPEGLSPPIGFAHQHYAKGNLVRICLGATI
                  230
                            240
                                    250
                                            260
                                                    270
   280
           290
                   300
                           310
                                   320
                                           330
inputs liligelæenhsrrkrlrhrgravorplpplpplpotrkshggodggrodvhsrglcs
     IIILIGILAEDWHSRKKCLQHRHRALQRPLPPLP-LA----
     280
             290
                     300
                            310
```

#### FIGURE 11

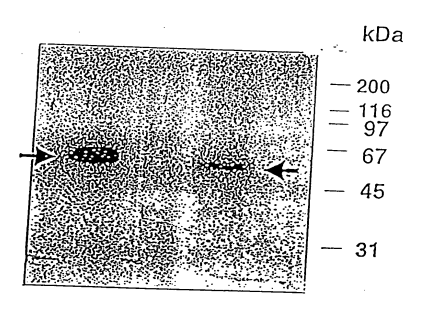


FIGURE 12

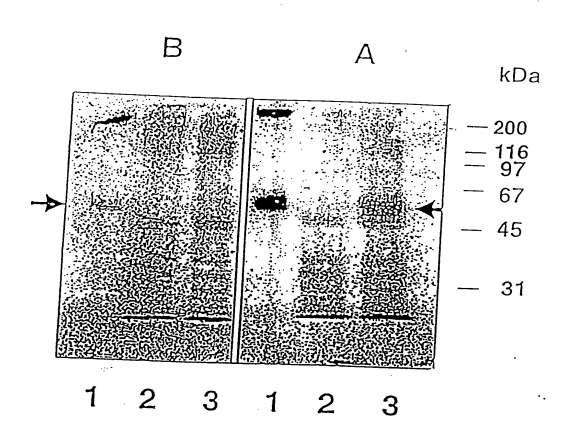


FIGURE 13

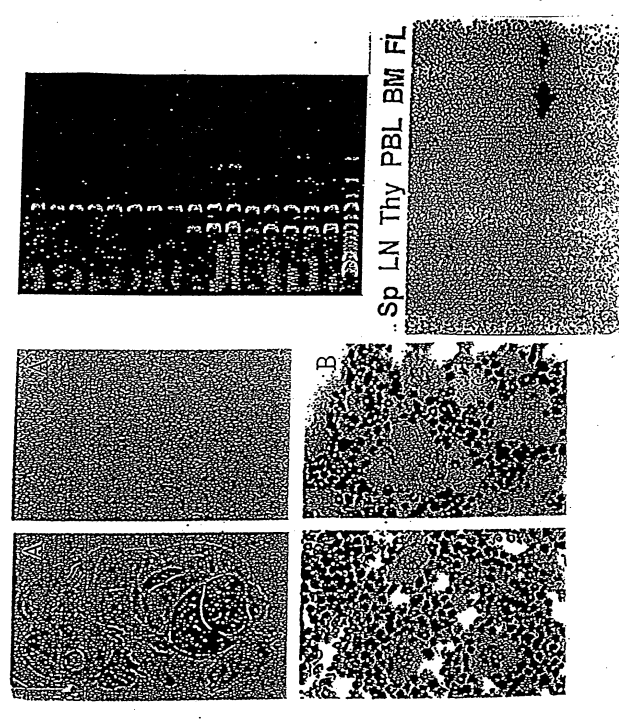
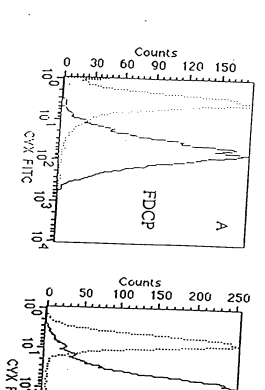
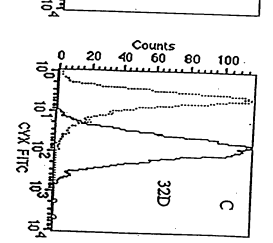


FIGURE 14





 $\boldsymbol{\omega}$ 

FIGURE 15

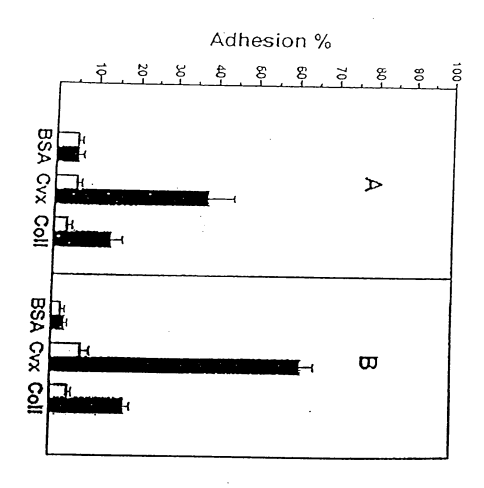


FIGURE 16

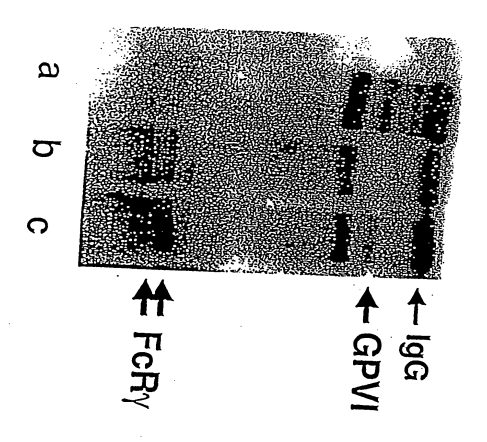
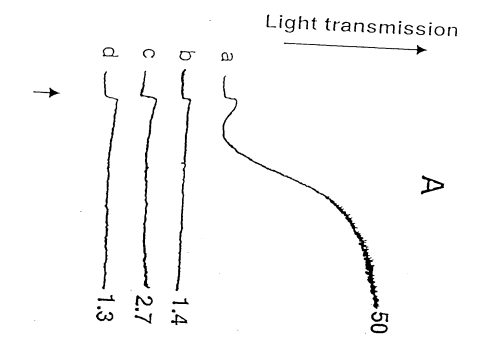


FIGURE 17



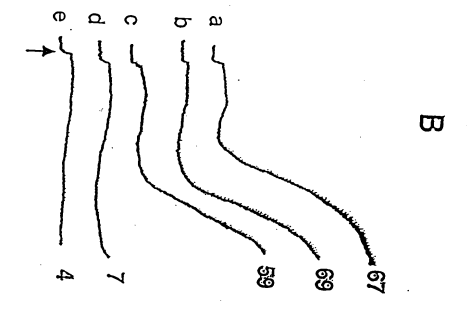


FIGURE 18

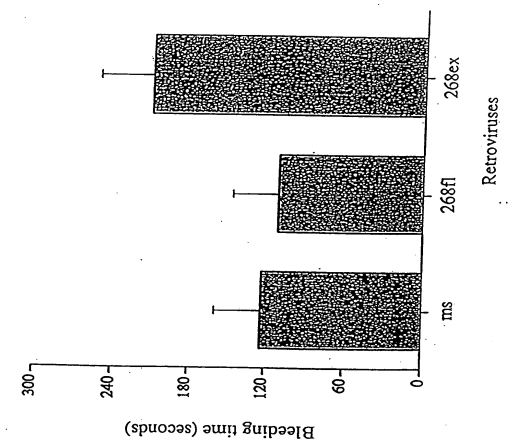


FIGURE 19

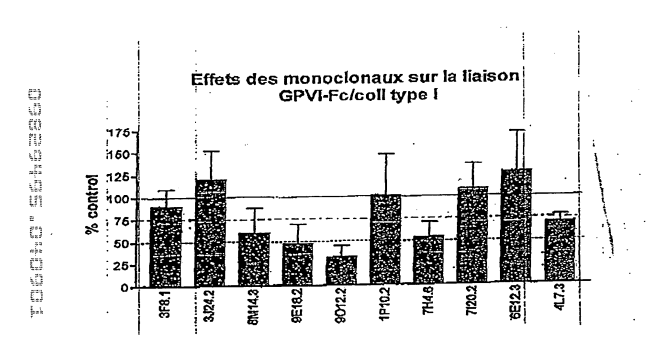
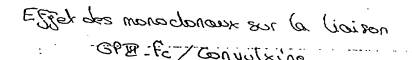


FIGURE 20



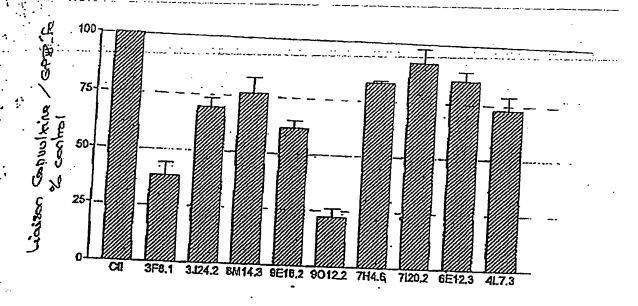
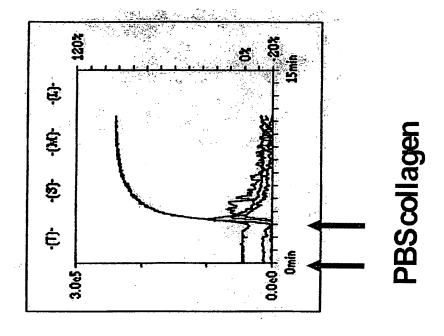


FIGURE 21



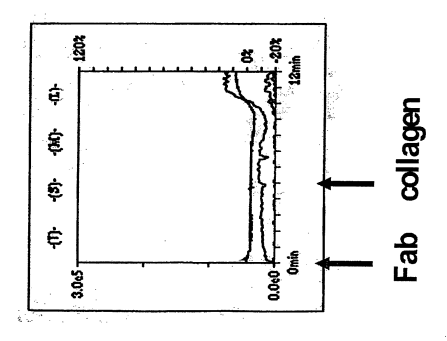


FIGURE 22

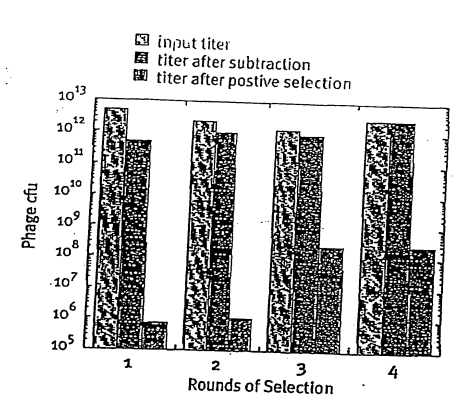


FIGURE 23

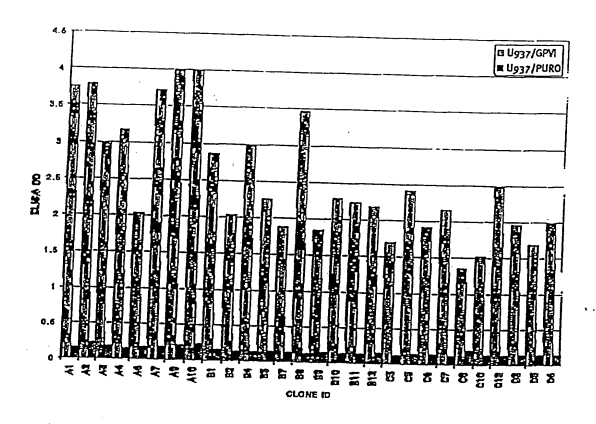


FIGURE 24a

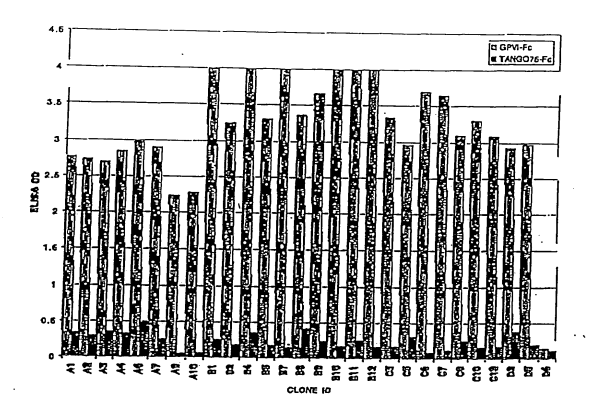


FIGURE 24b

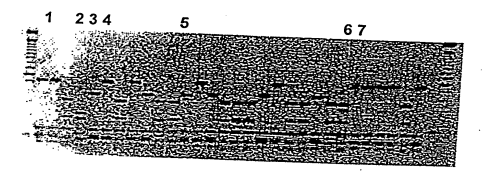


FIGURE 25

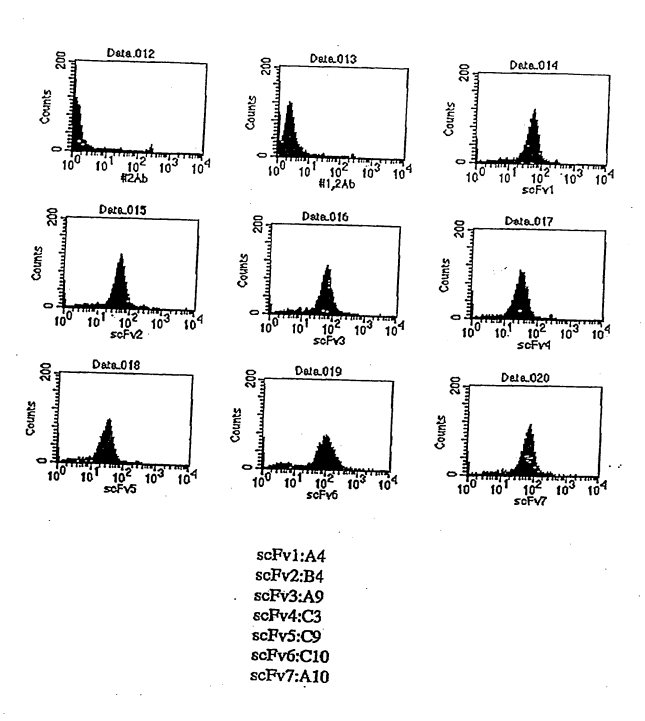


FIGURE 26

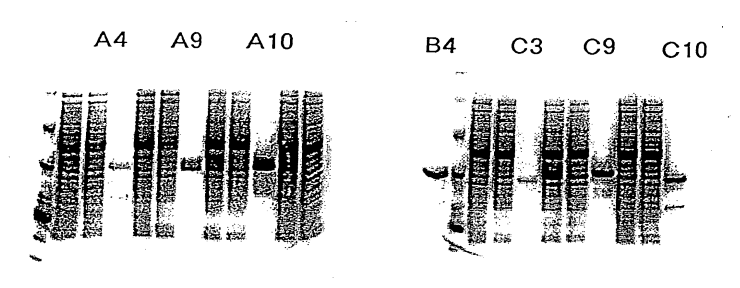


FIGURE 27

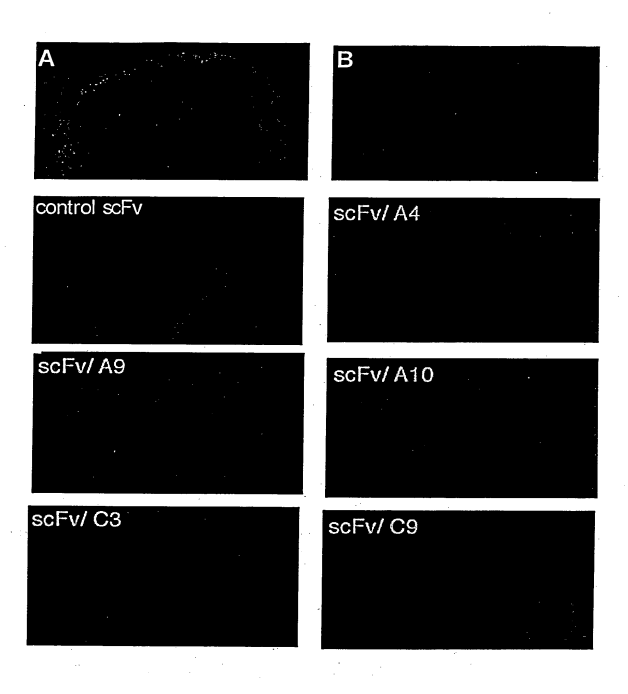


FIGURE 28

